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PCT

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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 5:	, ,	(11) International Publication Number:	WO 90/13653
C12N 15/62, C07K 13/00 C12P 21/02	A1	(43) International Publication Date:	15 November 1990 (15.11.90)
(21) International Application Number: PCT/GB (22) International Filing Date: 26 April 1990 ((30) Priority data: 8909916.2 29 April 1989 (29.04.89) (71) Applicant (for all designated States except US): DE OTECHNOLOGY LIMITED [GB/GB]; Cast Castle Boulevard, Nottingham NG7 1FD (GB)	(26.04.9 G ELTA B tle Cour	+ pean patent), CH (European tent), DK (European patent), FR (European patent), GB, IT (European patent), JP, NL (European patent), SE (B) Published With international search rep	n patent), DE (European pa-), ES (European patent). FI. GB (European patent). HU. KR, LU (European patent). European patenti. US.
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(57) Abstract

A fusion polypeptide comprising, as at least part of the N-terminal portion thereof, an N-terminal portion of HSA or a variant thereof and, as at least part of the C-terminal portion thereof, another polypeptide except that, when the said N-terminal portion of HSA is the 1-n portion where n is 369 to 419 or a variant thereof, then the said polypeptide is one of various specified entities, including the 585 to 1578 portion of human fibronectin or a variant thereof. The HSA-like portion may have additional N-terminal residues, such as secretion leader sequences (signal sequences). The C-terminal portion is preferably the 585-1578 portion of human plasma fibronectin. The N-terminal and C-terminal portions may be cleavable to yield the isolated C-terminal portion, with the N-terminal portion having served to facilitate secretion from the host.

One aspect of the present invention provides a fusion polypeptide comprising, as at least part of the N-terminal portion thereof, an N-terminal portion of HSA or a variant thereof and, as at least part of the C-terminal portion thereof, another polypeptide except that, when the said Nterminal portion of HSA is the 1-n portion where n is 369 to 419 or a variant thereof then the said polypeptide is (a) the 585 to 1578 portion of human fibronectin or a variant thereof, (b) the 1 to 368 portion of CD4 or a variant thereof, (c) platelet derived growth factor, or a variant thereof, (d) transforming growth factor, or a variant thereof, (e) the 1-261 portion of mature human plasma fibronectin or a variant thereof, (f) the 278-578 portion of mature human plasma fibronectin or a variant thereof, (g) the 1-272 portion of mature human von Willebrand's Pactor or a variant thereof, or (h) alpha-1-antitrypsin or a variant thereof.

The N-terminal portion of HSA is preferably the said 1-n portion, the 1-177 portion (up to and including the cysteine), the 1-200 portion (up to but excluding the cysteine) or a portion intermediate 1-177 and 1-200.

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The term "human serum albumin" (HSA) is intended to include (but not necessarily to be restricted to) known or yet-to-be-discovered polymorphic forms of HSA. example, albumin Naskapi has Lys-372 in place of Glu-372 and pro-albumin Christchurch has an altered pro-sequence. The term "variants" is intended to include (but not necessarily to be restricted to) minor artificial variations in sequence (such as molecules lacking one or a few residues, having conservative substitutions or minor insertions of residues, or having minor variations of amino acid structure). Thus polypeptides which have 80%, preferably 85%, 90%, 95% or 99%, homology with HSA are deemed to be "variants". It is also preferred for such variants to be physiologically equivalent to HSA; that is say, variants preferably share at least pharmacological utility with HSA. Furthermore, putative variant which is to be used pharmacologically should be non-immunogenic in the animal (especially human) being treated.

Conservative substitutions are those where one or more amino acids are substituted for others having similar properties such that one skilled in the art of polypeptide chemistry would expect at least the secondary structure, and preferably the tertiary structure, of the polypeptide to be substantially unchanged. For example, typical such

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substitutions include asparagine for glutamine, serine for asparagine and arginine for lysine. Variants may alternatively, or as well, lack up to ten (preferably only one or two) intermediate amino acid residues (ie not at the termini of the said N-terminal portion of HSA) in comparison with the corresponding portion of natural HSA; preferably any such omissions occur in the 100 to 369 portion of the molecule (relative to mature HSA itself) (if present). Similarly, up to ten, but preferably only one or two, amino acids may be added, again in the 100 to 369 portion for preference (if present). "physiologically functional equivalents" also encompasses larger molecules comprising the said sequence plus a further sequence at the N-terminal (for example, pro-HSA, pre-pro-HSA and met-HSA).

Clearly, the said "another polypeptide" in the fusion compounds of the invention cannot be the remaining portion of HSA, since otherwise the whole polypeptide would be HSA, which would not then be a "fusion polypeptide".

Even when the HSA-like portion is not the said 1-n portion of HSA, it is preferred for the non-HSA portion to be one of the said (a) to (h) entities.

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The 1 to 368 portion of CD4 represents the first four disulphide-linked immunoglobulin-like domains of the human T lymphocyte CD4 protein, the gene for and amino acid sequence of which are disclosed in D. Smith et al (1987) Science 328, 1704-1707. It is used to combat HIV infections.

The sequence of human platelet-derived growth factor (PDGF) is described in Collins et al (1985) Nature 316, 748-750. Similarly, the sequence of transforming growth factors β (TGF- β) is described in Derynck et al (1985) Nature 316, 701-705. These growth factors are useful for wound-healing.

A cDNA sequence for the 1-261 portion of Fn was disclosed in EP-A-207 751 (obtained from plasmid pFH6 with endonuclease PvuII). This portion binds fibrin and can be used to direct fused compounds to blood clots.

A cDNA sequence for the 278-578 portion of Fn, which contains a collagen-binding domain, was disclosed by R.J. Owens and F.E. Baralle in 1986 E.M.B.O.J. $\underline{5}$, 2825-2830. This portion will bind to platelets.

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The 1-272 portion of von Willebrand's Factor binds and stabilises factor VIII. The sequence is given in Bontham et al, Nucl. Acids Res. 14, 7125-7127.

Variants of alpha-1-antitrypsin include those disclosed by Rosenburg et al (1984) Nature 312, 77-80. In particular, the present invention includes the Pittsburgh variant (Met³⁵⁸ is mutated to Arg) and the variant where Pro³⁵⁷ and Met³⁵⁸ are mutated to alanine and arginine respectively. These compounds are useful in the treatment of septic shock and lung disorders.

Variants of the non-HSA portion of the polypeptides of the invention include variations as discussed above in relation to the HSA portion, including those with conservative amino acid substitutions, and also homologues from other species.

The fusion polypeptides of the invention may have N-terminal amino acids which extend beyond the portion corresponding to the N-terminal portion of HSA. For example, if the HSA-like portion corresponds to an N-terminal portion of mature HSA, then pre-, pro-, or pre-pro sequences may be added thereto, for example the yeast alpha-factor leader sequence. The fused leader portions of WO 90/01063 may be used. The polypeptide which is

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fused to the HSA portion may be a naturally-occurring polypeptide, a fragment thereof or a novel polypeptide, including a fusion polypeptide. For example, in Example 3 below, a fragment of fibronectin is fused to the HSA portion via a 4 amino acid linker.

It has been found that the amino terminal portion of the HSA molecule is so structured as to favour particularly efficient translocation and export of the fusion compounds of the invention in eukaryotic cells.

A second aspect of the invention provides a transformed host having a nucleotide sequence so arranged as to express a fusion polypeptide as described above. By "so arranged", we mean, for example, that the nucleotide sequence is in correct reading frame with an appropriate RNA polymerase binding site and translation start sequence and is under the control of a suitable promoter. The promoter may be homologous with or heterologous to the host. Downstream (3') regulatory sequences may be included if desired, as is known. The host is preferably yeast (for example Saccharomyces spp., e.g. S. cerevisiae; Kluyveromyces spp., e.g. K. lactis; Pichia spp.; or Schizosaccharomyces spp., e.g. S. pombe) but may be any

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other suitable host such as $\underline{E.\ coli}$, $\underline{B.\ subtilis}$, Aspergillus spp., mammalian cells, plant cells or insect cells.

A third aspect of the invention provides a process for preparing a fusion polypeptide according to the first aspect of the invention by cultivation of a transformed host according to the second aspect of the invention, followed by separation of the fusion polypeptide in a useful form.

A fourth aspect of the invention provides therapeutic methods of treatment of the human or other animal body comprising administration of such a fusion polypeptide.

In the methods of the invention we are particularly concerned to improve the efficiency of secretion of useful therapeutic human proteins from yeast and have conceived the idea of fusing to amino-terminal portions of HSA those proteins which may ordinarily be only inefficiently secreted. One such protein is a potentially valuable wound-healing polypeptide representing amino acids 585 to 1578 of human fibronectin (referred to herein as Fn 585-1578). As we have described in a separate application (filed simultaneously herewith) this molecule contains cell spreading, chemotactic and chemokinetic activities

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useful in healing wounds. The fusion polypeptides of the present invention wherein the C-terminal portion is Fn 585-1578 can be used for wound healing applications as biosynthesised, especially where the hybrid human protein will be topically applied. However, the portion representing amino acids 585 to 1578 of human fibronectin can if desired be recovered from the fusion protein by preceding the first amino acid of the fibronectin portion by amino acids comprising a factor X cleavage site. After isolation of the fusion protein from culture supernatant, the desired molecule is released by factor X cleavage and purified by suitable chromatography (e.g. ion-exchange chromatography). Other sites providing for enzymatic cr chemical cleavage can be provided, either by appropriate juxtaposition of the N-terminal and C-terminal portions or by the insertion therebetween of an appropriate linker.

At least some of the fusion polypeptides of the invention, especially those including the said CD4 and vWF fragments, PDGF and $\alpha_1 AT$, also have an increased half-life in the blood and therefore have advantages and therapeutic utilities themselves, namely the therapeutic utility of the non-HSA portion of the molecule. In the case of $\alpha_1 AT$ and others, the compound will normally be administered as

a one-off dose or only a few doses over a short period, rather than over a long period, and therefore the compounds are less likely to cause an immune response.

EXAMPLES : SUMMARY

Standard recombinant DNA procedures were as described by Maniatis et al (1982 and recent 2nd edition) unless otherwise stated. Construction and analysis of phage M13 recombinant clones was as described by Messing (1983) and Sanger et al (1977).

DNA sequences encoding portions of human serum albumin used in the construction of the following molecules are derived from the plasmids mHOB12 and pDBD2 (EP-A-322 094, Delta Biotechnology Ltd, relevant portions of which are reproduced below) or by synthesis of oligonucleotides equivalent to parts of this sequence. DNA sequences encoding portions of human fibronectin are derived from the plasmid pFHDEL1, or by synthesis of oligonucleotides equivalent to parts of this sequence. Plasmid pFHDEL1, which contains the complete human cDNA encoding plasma fibronectin, was obtained by ligation of DNA derived from plasmids pFH6, 16, 54, 154 and 1 (EP-A-207 751; Delta Biotechnology Ltd).

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This DNA represents an mRNA variant which does not contain the 'ED' sequence and had an 89-amino acid variant of the III-CS region (R.J. Owens, A.R. Kornblihtt and F.E. Baralle (1986) Oxford Surveys on Eukaryotic Genes 3 141-160). The map of this vector is disclosed in Fig. 11 and the protein sequence of the mature polypeptide produced by expression of this cDNA is shown in Fig. 5.

Oligonucleotides were synthesised on an Applied Biosystems 380B oligonucleotide synthesiser according to the manufacturer's recommendations (Applied Biosystems, Warrington, Cheshire, UK).

An expression vector was constructed in which DNA encoding the HSA secretion signal and mature HSA up to and including the 387th amino acid, leucine, fused in frame to DNA encoding a segment of human fibronectin representing amino acids 585 to 1578 inclusive, was placed downstream of the hybrid promoter of EP-A-258 067 (Delta Biotechnology), which is a highly efficient galactose-inducible promoter functional in Saccharomyces cerevisiae. The codon for the 1578th amino acid of human fibronectin was directly followed by a stop codon (TAA) and then the S. cerevisiae phosphoglycerate kinase (PGK) gene transcription terminator. This vector was then introduced into S. cerevisiae by transformation, wherein it directed

the expression and secretion from the cells of a hybrid molecule representing the N-terminal 387 amino acids of HSA C-terminally fused to amino acids 585 to 1578 of human fibronectin.

In a second example a similar vector is constructed so as to enable secretion by <u>S. cerevisiae</u> of a hybrid molecule representing the N-terminal 195 amino acids of HSA C-terminally fused to amino acids 585 to 1578 of human fibronectin.

Aspects of the present invention will now be described by way of example and with reference to the accompanying drawings, in which:

Figure 1 (on two sheets) depicts the amino acid sequence currently thought to be the most representative of natural HSA, with (boxed) the alternative C-termini of HSA(1-n);

Figure 2 (on two sheets) depicts the DNA sequence coding for mature HSA, wherein the sequence included in Linker 3 is underlined;

Figure 3 illustrates, diagrammatically, the construction of mHOB16;

Figure 4 illustrates, diagrammatically, the construction of pHOE31;

Figure 5 (on 6 sheets) illustrates the mature protein sequence encoded by the Fn plasmid pFHDEL1;

Figure 6 illustrates Linker 5, showing the eight constituent oligonucleotides;

Figure 7 shows schematically the construction of plasmid pDBDF2;

Figure 8 shows schematically the construction of plasmid pDBDF5;

Figure 9 shows schematically the construction of plasmid pDBDF9;

Figure 10 shows schematically the construction of plasmid DBDF12, using plasmid pFHDEL1; and

Figure 11 shows a map of plasmid pFHDEL1.

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EXAMPLE 1 : HSA 1-387 FUSED TO Fn 585-1578

The following is an account of a preparation of plasmids comprising sequences encoding a portion of HSA, as is disclosed in EP-A-322 094.

The human serum albumin coding sequence used in the construction of the following molecules is derived from the plasmid M13mp19.7 (EP-A-201 239, Delta Biotech- nology Ltd.) or by synthesis of oligonucleotides equivalent to parts of this sequence. Oligonucleotides were synthesised using phosphoramidite chemistry on an Applied Biosystems 380B oligonucleotide synthesizer according to the manufacturer's recommendations (AB Inc., Warrington, Cheshire, England).

An oligonucleotide was synthesised (Linker A) which represented a part of the known HSA coding sequence (Figure 2) from the PstI site (1235-1240, Figure 2) to the codon for valine 381 wherein that codon was changed from GTG to GTC:

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Li	nl	ke	I	1

	D ·	P	H	E	С	Ā			
5′	GAT	CCT	CAT	GAA	TGC	TAT			
3' ACGT	CTA	GGA	GTA	CTT	ACG	ATA			
1247									

A	K	v	F	D	E	F	K
GCC	AAA .	GTG	TTC	GAT	GAA	TTT	AAA
CGG	TTT	CAC	AAG	CTA	CTT	AAA	TTT
		1267					

P L V
CTT GTC 3'
GGA CAG 5'

Linker 1 was ligated into the vector M13mp19 (Norrander et al, 1983) which had been digested with PstI and HincII and the ligation mixture was used to transfect E.coli strain XL1-Blue (Stratagene Cloning Systems, San Diego, CA). Recombinant clones were identified by their failure to evolve a blue colour on medium containing the chromogenic indicator X-gal (5-bromo-4-chloro-3-indolyl-β-D-galactoside) in the present of IPTG (isopropylthio-β-galactoside). DNA sequence analysis of template DNA prepared from bacteriophage particles of recombinant clones identified a molecule with the required DNA sequence, designated mHOB12 (Figure 3).

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M13mpl9.7 consists of the coding region of mature HSA in M13mpl9 (Norrander et al, 1983) such that the codon for the first amino acid of HSA, GAT, overlaps a unique $\underline{\text{Xho}}$ I site thus:

Asp Ala

- 5' CTCGAGATGCA 3'
- 3' GAGCTCTACGT 5'

<u>Xho</u>I

(EP-A-210 239). M13mp19.7 was digested with XhoI and made flush-ended by S1-nuclease treatment and was then ligated with the following oligonucleotide (Linker 2):

Linker 2

5' T C T T T T A T C C A A G C T T G G A T A A A A G A 3'
3' A G A A A T A G G T T C G A A C C T A T T T T C T 5'

<u>Hin</u>dIII

The ligation mix was then used to transfect <u>E.coli</u> XL1-Blue and template DNA was prepared from several plaques and then analysed by DNA sequencing to identify a clone, pDBD1 (Figure 4), with the correct sequence.

A 1.1 kb HindIII to PstI fragment representing the 5' end of the HSA coding region and one half of the inserted oligonucleotide linker was isolated from pDBD1 by agarose gel electrophoresis. This fragment was then ligated with double stranded mHOB12 previously digested with HindIII and PstI and the ligation mix was then used to transfect Single stranded template DNA was E.coli XL1-Blue. prepared from mature bacteriophage particles of several plaques. The DNA was made double stranded in vitro by extension from annealed sequencing primer with the Klenow fragment of DNA polymerase I in the presence of decxynucleoside triphosphates. Restriction enzyme analysis of this DNA permitted the identification of a clone with the correct configuration, mHOB15 (Figure 4).

The following oligonucleotide (Linker 3) represents from the codon for the 382nd amino acid of mature HSA (glutamate, GAA) to the codon for lysine 389 which is followed by a stop codon (TAA) and a <u>HindIII</u> site and then a BamHI cohesive end:

Linker 3

- E E P O N L I K J
- 5' GAA GAG CCT CAG AAT TTA ATC AAA TAA GCTTG 3'
- 3' CTT CTC GGA GTC TTA AAT TAG TTT ATT CGAACCTAG 5'

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This was ligated into double stranded mHOB15, previously digested with <u>HincII</u> and <u>BamHI</u>. After ligation, the DNA was digested with <u>HincII</u> to destroy all non-recombinant molecules and then used to transfect <u>E.coli</u> <u>KL1-Blue</u>. Single stranded DNA was prepared from bacteriophage particles of a number of clones and subjected to DNA sequence analysis. One clone having the correct DNA sequence was designated mHOB16 (Figure 4).

A molecule in which the mature HSA coding region was fused to the HSA secretion signal was created by insertion of Linker 4 into <u>BamHI</u> and <a href="mailto:XhoI digested Ml3mpl9.7 to form pDBD2 (Figure 4).

Linker 4

I

ATT

TAA

s

TCC

AGG

L

CTT

GAA

	M	K	W	v	S	F
5' GATCC	ATG	AAG	TGG	GTA	AGC	TTT
G	TAC	TTC	ACC	CAT	TCG	AAA
	-					

L

CTT

GAA

F

TTT

AAA

L

CTC

GAG AAA

s

AGC

TCG

F

TTT

19 .

S Α Y S R G TCG GCT TAT TCC AGG GGT GTG Tar. AGC CGA ATA AGG TCC CCA CAC AAA

R R CG 3'

In this linker the codon for the fourth amino acid after the initial methionine, ACC for threonine in the HSA prepro leader sequence (Lawn et al, 1981), has been changed to AGC for serine to create a HindIII site.

A sequence of synthetic DNA representing a part of the known HSA coding sequence (Lawn et al., 1981) (amino acids 382 to 387, Fig. 2), fused to part of the known fibronectin coding sequence (Kornblihtt et al., 1985) (amino acids 585 to 640, Fig. 2), was prepared by synthesising six oligonucleotides (Linker 5, Fig. 6). The oligonucleotides 2, 3, 4, 6, 7 and 8 were phosphorylated polynucleotide the kinase and then using T4oligonucleotides were annealed under standard conditions in pairs, i.e. 1+8, 2+7, 3+6 and 4+5. The annealed oligonucleotides were then mixed together and ligated with mHOB12 which had previously been digested with the restriction enzymes HincII and EcoRI. The ligation

mixture was then used to transfect <u>E.coli</u> XL1-Blue (Stratagene Cloning Systems, San Diego, CA). Single stranded template DNA was then prepared from mature bacteriophage particles derived from several independent plaques and then was analysed by DNA sequencing. A clone in which a linker of the expected sequence had been correctly inserted into the vector was designated pDBDF1 (Fig. 7). This plasmid was then digested with <u>PstI</u> and <u>EcoRI</u> and the approx. 0.24kb fragment was purified and then ligated with the 1.29kb <u>BamHI-PstI</u> fragment of pDBD2 (Fig. 7) and <u>BamHI + EcoRI</u> digested pUC19 (Yanisch-Perron, et al., 1985) to form pDBDF2 (Fig. 7).

A plasmid containing a DNA sequence encoding full length human fibronectin, pFHDEL1, was digested with <u>EcoRI</u> and <u>XhoI</u> and a 0.77kb <u>EcoRI-XhoI</u> fragment (Fig. 8) was isolated and then ligated with <u>EcoRI</u> and <u>SalI</u> digested M13 mp18 (Norrander et al., 1983) to form pDBDF3 (Fig. 8).

The following oligonucleotide linker (Linker 6) was synthesised, representing from the PstI site at 4784-4791 of the fibronectin sequence of EP-A-207 751 to the codon for tyrosine 1578 (Fig. 5) which is followed by a stop codon (TAA), a HindIII site and then a BamHI cohesive end:

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Linker 6

 ${\sf G}$ P D Q T E M T I E G L ${\sf GGT} \ \ {\sf CCA} \ \ {\sf GAT} \ \ {\sf CAA} \ \ {\sf ACA} \ \ {\sf GAA} \ \ {\sf ATG} \ \ {\sf ACT} \ \ {\sf ATT} \ \ {\sf GAA} \ \ {\sf GGC} \ \ {\sf TTG}$ A CGT CCA GGT CTA GTT TGT CTT TAC TGA TAA CTT CCG AAC

Q P T V E Y Stop

CAG CCC ACA GTG GAG TAT TAA GCTTG

GTC GGG TGT CAC CTC ATA ATT CGAACCTAG

This linker was then ligated with PStI and HindIII digested pDBDF3 to form pDBDF4 (Fig. 8). The following DNA fragments were then ligated together with BclII digested pKV50 (EP-A-258 067) as shown in Fig. 8: 0.68kb ECORI-BamHI fragment of pDBDF4, 1.5kb BamHI-StuI fragment of pDBDF2 and the 2.2kb StuI-ECORI fragment of pFHDEL1. The resultant plasmid pDBDF5 (Fig. 8) includes the promoter of EP-A-258 067 to direct the expression of the HSA secretion signal fused to DNA encoding amino acids 1-387 of mature HSA, in turn fused directly and in frame with DNA encoding amino acids 585-1578 of human fibronectin, after which translation would terminate at the stop codon TAA. This is then followed by the S.cerevisiae PGK gene transcription terminator. The

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plasmid also contains sequences which permit selection and maintenance in <u>Escherichia coli</u> and <u>S.cerevisiae</u> (EP-A-258 067).

This plasmid was introduced into <u>S.cerevisiae</u> S150-2E (<u>leu2-3 leu2-112 ura3-52 trpl-289 his3- 1</u>) by standard procedures (Beggs, 1978). Transformants were subsequently analysed and found to produce the HSA-fibronectin fusion protein.

EXAMPLE 2 : HSA 1-195 FUSED TO Fn 585-1578

In this second example the first domain of human serum albumin (amino acids 1-195) is fused to amino acids 585-1578 of human fibronectin.

The plasmid pDBD2 was digested with BamHI and BclII and the 0.79kb fragment was purified and then ligated with BamHI-digested M13mp19 to form pDBDF6 (Fig. 6). The following oligonucleotide:

5'-C C A A A G C T C G A G G A A C T T C G-3'

was used as a mutagenic primer to create a <u>Xho</u>I site in pDBDF6 by <u>in vitro</u> mutagenesis using a kit supplied by Amersham International PLC. This site was created by

changing base number 696 of HSA from a T to a G (Fig. 2). The plasmid thus formed was designated pDBDF7 (Fig. 9). The following linker was then synthesised to represent from this newly created XhoI site to the codon for lysine 195 of HSA (AAA) and then from the codon for isoleucine 585 of fibronectin to the ends of oligonucleotides 1 and 8 shown in Fig. 6.

Linker 7

D E L R D E G K A S S A K

TC GAT GAA CTT CGG GAT GAA GGG AAG GCT TCG TCT GCC AAA

A CTT GAA GCC CTA CTT CCC TTC CGA AGC AGA CGG TTT

I T E T P S Q P N S H

ATC ACT GAG ACT CCG AGT CAG C

TAG TGA CTC TGA GGC TCA GTC GGG TTG AGG GTG G

This linker was ligated with the annealed oligonucleotides shown in Fig. 3, i.e. 2+7, 3+6 and 4+5 together with XhoI and EcoRI digested pDBDF7 to form pDBDF8 (Fig. 9). Note that in order to recreate the original HSA DNA sequence, and hence amino acid sequence, insertion of linker 7 and the other oligonucleotides into pDBDF7 does not recreate the XhoI site.

The 0.83kb <u>BamHI-StuI</u> fragment of pDBDF8 was purified and then was ligated with the 0.68kb <u>EcoRI-BamHI</u> fragment of pDBDF2 and the 2.22kb <u>StuI-EcoRI</u> fragment of pFKDEL1 into <u>BqlII-digested</u> pKV50 to form pDBDF9 (Fig. 9). This plasmid is similar to pDBDF5 except that it specifies only residues 1-195 of HSA rather than 1-387 as in pDBDF5.

When introduced into <u>S.cerevisiae</u> S150-2B as above, the plasmid directed the expression and secretion of a hybrid molecule composed of residues 1-195 of HSA fused to residues 585-1578 of fibronectin.

EXAMPLE 3 : HSA 1-387 FUSED TO Fn 585-1578, AS CLEAVABLE MOLECULE

In order to facilitate production of large amounts of residues 585-1578 of fibronectin, a construct was made in which DNA encoding residues 1-387 of HSA was separated from DNA encoding residues 585-1578 of fibronectin by the sequence

I E G R
ATT GAA GGT AGA
TAA CTT CCA TCT

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which specifies the cleavage recognition site for the blood clotting Factor X. Consequently the purified secreted product can be treated with Factor X and then the fibronectin part of the molecule can be separated from the HSA part.

To do this two oligonucleotides were synthesised and then annealed to form Linker 8.

Linker 8

E E P Q N L I E G

GAA GAG CCT CAG AAT TTA ATT GAA GGT

CTT CTC GGA GTC TTA AAT TAA CTT CCA

R I T E T P S Q P

AGA ATC ACT GAG ACT CCG AGT CAG C

TCT TAG TGA CTC TGA GGC TCA GTC GGG

N S H

TTG AGG GTG G

This linker was then ligated with the annealed oligonucleotides shown in Fig. 6, i.e. 2+7, 3+6 and 4+5 into <u>Hin</u>cII and <u>Eco</u>RI digested mHOB12, to form pDEDF10

(Fig. 7). The plasmid was then digested with <u>PstI</u> and <u>EcoRI</u> and the roughly 0.24kb fragment was purified and then ligated with the 1.29kb <u>BamHI-PstI</u> fragment of pDBD2 and <u>BamHI</u> and <u>EcoRI</u> digested pUCl9 to form pDBDF11 (Fig. 10).

The 1.5kb BamHI-StuI fragment of pDBDF11 was then ligated with the 0.68kb EcoRI-BamH1 fragment of pDBDF4 and the 2.22kb StuI-EcoRI fragment of pFHDEL1 into BclII-digested pKV50 to form pDBDF12 (Fig. 10). This plasmid was then introduced into S.cerevisiae S150-2B. The purified secreted fusion protein was treated with Factor X to liberate the fibronectin fragment representing residues 585-1578 of the native molecule.

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CLAIMS

A fusion polypeptide comprising, as at least part of the N-terminal portion thereof, an N-terminal portion of HSA or a variant thereof and, as at least part of the C-terminal portion thereof, another polypeptide except that, when the said N-terminal portion of HSA is the 1-n portion where n is 369 to 419 or a variant thereof then the said polypeptide is (a) the 585 to 1578 portion of human fibronectin or a variant thereof, (b) the 1 to 368 portion of CD4 or a variant thereof, (c) platelet derived growth factor or a variant thereof, (d) transforming growth factor β or a variant thereof, (e) the 1-261 portion of mature human plasma fibronectin or a variant thereof, (f) 278-578 portion of mature human plasma fibronectin or a variant thereof, (g) the 1-272 portion of mature human von Willebrand's Factor or a variant thereof, or (h) alpha-l-antitrypsin or a variant thereof.

- A fusion polypeptide according to Claim 1
 additionally comprising at least one N-terminal amino
 acid extending beyond the portion corresponding to
 the N-terminal portion of HSA.
- 3. A fusion polypeptide according to Claim 1 or 2 wherein there is a cleavable region at the junction of the said N-terminal or C-terminal portions.
- 4. A fusion polypeptide according to any one of the preceding claims wherein the said C-terminal portion is the 585 to 1578 portion of human plasma fibronectin or a variant thereof.
- 5. A transformed or transfected host having a nucleotide sequence so arranged as to express a fusion polypeptide according to any one of the preceding claims.
- 6. A process for preparing a fusion polypeptide by cultivation of a host according to Claim 5, followed by separation of the fusion polypeptide in a useful form.
- A fusion polypeptide according to any one of Claims 1 to 4 for use in therapy.

FIGURE_1

λs	al.	a H1.	s Ly:	s 5e.	- 5 1:	: Va.	ندا	a Hi	1 جہ ہ		e Ly:	s As	p Le	(G1 د	y 31	u 51	u As	n Pn	20 e Lys
Ala	ı Le	ı Va.	l le:	2 Il	e Ala	. Phe	e Al	e G1:	3: : "Y		: Gl:	n Gl	n Cys	; P r :	o Ph	e Gi	u As	p X1	40 15 Vai
Lys	i Let	ı Val	l Asi	n Gl:	. Val	Tin.	Gli	ı Phe	50 114 e	-	: Thi	Су:	s Val	. Ala	As:	p Gl:	: Se:	- Ala	50 2 Glu
Ast	. Cys	s As;	p Lys	5 5e:	: Leu	<u> </u>		- Leu	70 Phe		, ys i) Lys	s Leu	Cys	: Thi	va!	. Ala	Th	80 Leu
Arç	: 61:		Tyz	- Gl;	/ Glu	. Met	. Ala	a Asț	90 Cys		Ala	Lys	Gla	Glu	. Pr:	: G1:	: Arq	Ast	100 1012
Cys	Phe	. Leu	: Glm	: His	: Lys	Asp	λs) Asn	110		Leu	Pro) Arç	Leu	. Val	. Arg	270	glu	120 Val
ASP	Vai	. Met	Cys	The	· Ala	Phe	His	ASP	130 Asn		Glu	<u> </u>	Phe	Leu	Lys	: Lys	Tyr	Leu	140 Ty=
Glu	Ile	Ala	Arg	YIS	Hls	Pro	Tys	Phe	150 Ty=		Pro	Glu	Leu	Leu	Ph∈	Pie	Ala	Lys	160 کتو
Tyr	Lys	Ala	Ala	Phe		Şlu	Cys	Cys	170 Gln		Als	λsp	Lys	àla	Ala	Cys	Leu	Leu	:80
Lys	Leu	λsp	Glu	Leu	Arş	ĄSP.	Glu	Gly	190 Lys	Αlz	Ser	Ser	Ala	Lys	Glm	Arş	Leu	Lys	200 Cys
Ala	Ser	Leu	Gla	Lys	Phe	Gly	Glu	Arg.	210 Ala	Phe	Lys	Ala	?rp	Ala	Val	Ala	Arş	Leu	220 Ser
۵ln	λIŞ	Phe	250	Lys	Ala	510	Phe	Ala	230 Glu	Val	Ser	Lys	Leu	Val	71.	Asp	Leu	The	240 Lys
Val	Eis	7:-	Glu	Cys,	Cys	His	Gly	γs⊃	250 Lau	Leu	Glu	Cys	Ala	Asp	ÇZÁ	Мâ	Ala	Asp	250 Leú
Alz	Lys	Tyr	Ile	Cys	Glu	Asn	Gln	ςεk	270 Ser	Ile	Ser	Ser	Lys	Leu	Lys	Glu	Cys	Cys	280 Glu
Lys	?ro	Leu	Leu	Glu	Lys	5 er	Eis	Cys	290 11e	Ala	Glu	Val	Glu	Asn	Ąsp	Slu	Met	Pro	300 %12
Asp	Leu	250	Ser	Leu	Ala	lla	ζZK	Phe	310 Val	Glu	Ser	Lys	ÇZÁ	Val	Cy⁄s	Lys	ÀSTI	Tyr	320 Ala
Gī₽	Ala	Lys	çzƙ	Val	Phe	leu	Sly		330 Phe	Leu	Tyr	2]2	Tys	Ala	AIŞ	λī;	His	Pro	340 Asp
Tyr	Ser	Val	Vai	Leu	Lau	Lau	Arş		350 Ale	lys	Th <u>-</u> '	Tyr	Glu '	The -	7.5 <i>=</i>	Leu	51::		350 Cys
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FIGURE 1 Com	<u></u>		
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		430	440
Pro The Lau Va	al Glu Val Ser Arg Aso	Leu Gly Lys Val Gly Ser	Lys Cys Cys Lys His
		450	450
Pro Glu Ala Ly	vs Arq Met Pro Cys Ala	Glu Asp Tyr Leu Ser Val	Val leu Asm Glm Leu
·			
		470	450
Cvs Val Leu Hi	is Glu Lys Thr Pro Val	Ser Asp Arg Val Thr Lys	Cvs Cvs The Glu Ser
•	·		
		490	500
leu Val Asm Ar	eg Arg Pro Cys-Phe Ser	Ala Leu Glu Val Asp Glu	The Twe Val Pro Lys
		•	•
		510	520
Glu, Phe Asm Al	a Glu Thr Phe Thr Phe	His Ala Asp Ile Cvs Thr	Leu Ser Glu Lys Glu
•			
		530	540
Ard Gln Ile Ly:	s Lys Gln Thr Ala Leu	Val Glu Leu Val Lys His	Lvs Pro Lys Ala Thr
•	•	·	
	•	550	560
Lvs Glu Gla Le	er Lvs Ala Val Met Aso	Asp Phe Ala Ala Phe Val	Glu Lvs Cvs Cys Lys
5 , 5 522 523 25		•	
		570	580
Ala ASD ASD Eve	s Glu Thr Cys Phe Ala	Glu Glu Gly Lys Lys Leu	val Ala Ala Ser Gim
	, 		
Ale Ale Leu Gly	ນ ໂລນ		
, see or or .			

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FIGURE 2 DNA sequence coding for mature HSA

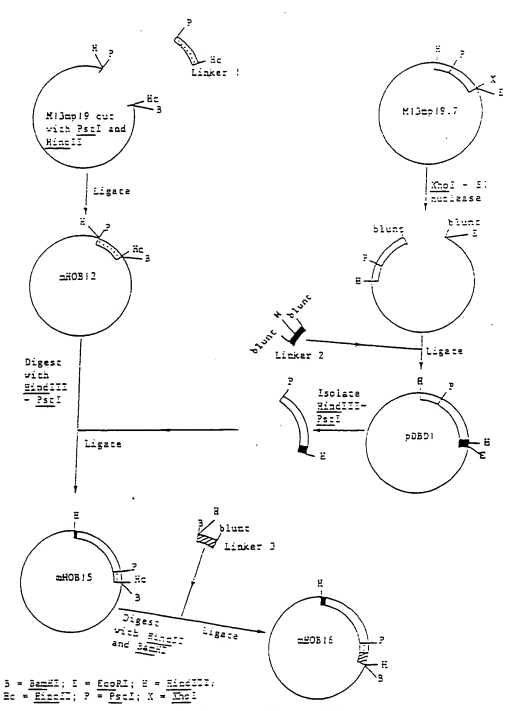
10	20	30	40	50	60	78	8.0
GATGCACACAAGAG D A H K S	TGAGGTTGCT E V A	CATCSGTTI H R F	X D L	G E E N	TTCAAAGCCT F X A	TGGTGTTGAT	A F
				130			
TGCTCAGTATCTTC	AGCAGTGTCC O O C P	ATTTGAAGA F E D	TCATGTAAA; X V E	ATTAGTGAATGA L V N E	AGTAACTGAA : V T E	TTTGCAAAAA 7 A K	CATGIG I C
170				210			
TTGCTGATGAGTCA	SCTGÄÄÄÄÄTT	GTGACAAAT	CACTTCATAC	CCTTTTTGGAG	ACAAATTATG	CACAGTTGCA	ACTOTT
V A D E 5	A E N	C D K .	SLHT	L F G	ם א ב כ	t v A	= =
250 CGTGAAACCTATGGT	260			290		310	
R E T Y G	E X A	D C C	y K Ö	E PER	N E C	F 1 0 H	Z Z
230	340	350	360	370	380	390	400
TGACAACCCAAACCT D N P N L							
410 TTTTGAAAAATACT	420 Tatatgaaa:			450 - CTTTTATGCCCC			
F L K K Y							
				530			
TATAAAGCTGCTTTT Y K A A F							
570	580	590	600	610	520	630	640
TGAAGGGAAGGCTTC	GTCTGCCAAA	CAGAGACTO	AAATGTGCCA	GTCTCCAAAAA	TTTGGAGAAA	GAGETTTEAA	AGCAT
E G K A S	SAX	Q R L	KCA	SLQX	FGZ	E A F K	A
650 GGGCAGTGGCTCGCCT			680			710	720
W A V A R I							
	740			770			500
GTCCACACGGAATGCT	GCCATGGAG	TCTGCTTG	ATGTGCTGA	TGACAGGGGGG	CCTTGCCAA	TATATETGTO	بمنتد
• •							
810 TCAGGATTCGATCTCC	820	830	840	E50	560	570 Catteresaa	
Q D S I S	S X L	X Z C	C E K	P L L E	к 5 н С	: I A E	v
890	900	910	920	.930	940	950	960
AAAATGATGAGATGCC E N D E M P	TGCTGACTTG A D L	P S L	GCTGCTGATS A A D	TTTGTTGAAAGT F V Z S	XAGGATGTTT K D V	GCAAAAACTA C X N Y	TGCT A
97 0	980	990	1000	1010	1020	:030	1040
GAGGCAAAGGATGTCT:	TCCTGGGCAT F L G M	STITITGIA F L Y	TGAATATGCA E Y A	AGAAGGCATCC R R R P	TOATTACTCT S Y C	eregreered: 7 7 L :	rge r : :

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FIGURE 2 Cont. 1050 1060 1070 * * * * 2 GAGACTTGCCAAGACATATGAAAAÇCACTCTAGAGAAGTGCTGTGCCGCCGCAGAGCACATGAAGCCTATGAAACGCCAAAGTGC 1140 - 1150 :180 ::90 F D E F K P L V E E F Q N L E K Q N C E L F E C L G E :280 TACAAATTCCAGAATGEGGTATTAGTTCGTTACACCAAGAAGTACCCCAAGTGTCAACTCCAACTCTTGTAGAGGTCTC Y K F Q N A L L V R Y T K K V P Q V S T P T L V E V S 1350 1360 1310 1320 AAGAAACCTAGGAAAAGTGGGCCAGCAAATGTTGTAAACATCCTGAAGCAAAAAGAATGCCCTGTGCAGAAGAACATATCTAT R N L G K V G S K C C K H P E A K R M P C A E D Y L 1420 1430 $\verb"costggtcctgaaccagtatgtgtgcatgaagaaaacgccagtaagtgacagagtaactacaaaatgtgcacagagtgc$ S V V L N Q L C V L H E K T F V S D R V T K C C T E S TTGGTGAACAGGGGACGATGCTTTTCAGGTCTGGAAGTGGAAACATAGGTTCGGAAAGAGTTTAATGCTGAAACATT AACACAAGCCCAAGGCAACAAAAGAGCAACTGAAAGCTGTTATGGATGATTTCGCAGCTTTTTGTAGAGAAGTGCTGCAAG K H K P K A T K E Q L K A V M D D F A A F V E K C C K :720 GCTGACGATAAGGAGACCTGCTTTGCCGAGGAGGGTAAAAAACTTGTTGCTGCAAGTCAAGCTGCCTTAGGCTTATAACA A D D K E T C F A E E G K K L V A A S Q A A L G L

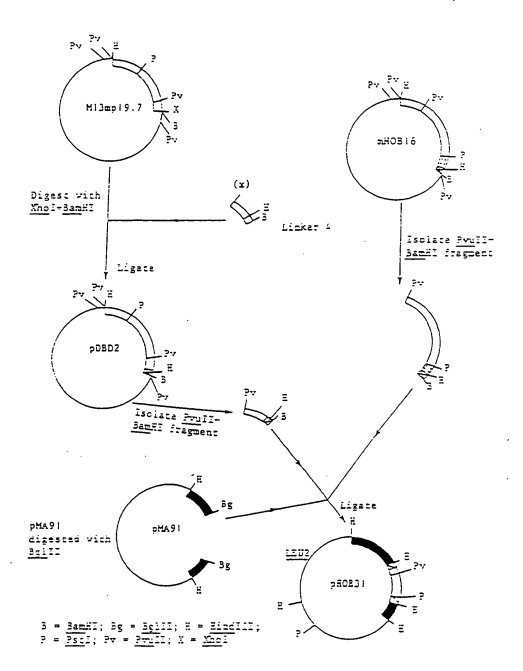
TOTACATTTAAAAGCATOTCAG

FIGURE 3 Construction of mHOBi6



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FIGURE 4 Construction of pHOB31



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Fig. 5A

900 00 00 00 00 00 00 00 200 220 Asn 180 80 Arg Asp Lys Ala Cys Arg n L Ş Asp Leu Met Arg 부 Ħ Ser GIN Thr Thr Thr Gln GIn Thr Lys 훋 ζŞ Asp HIS Trp Met Trp Lys Lys s Тyг Gly Arg Asn Lys Lys Asn Gly Arg **Asn His** Ser G S Asp Val Cys Val Asn <u>ช</u> Ala Ë Vα Leu Val Gin Ser <u>√</u> Ϋ́ Arg GIY cys Cys Phe S D ΕIS GΙ Ala Vai Asn GΙγ Ser 본 Gin Gin Trp Glu Arg Ser Ser GIN GIY 75 Trp ᇤ Ϋ́ ςλs Τŗ Phe Asn Cys 11e Ě <u>م</u> G Z His Leu Trp Cys Asn Met Lys Trp Cys Gly Asp HIS Thr Val Ser Phe Pro Phe Leu Tyr Thr Gly Asn Thr Met Leu Glu Lys Cys 뉴 Cys Gin Glu Thr Leu Pro Phe Thr 부 G S Asn Gly Arg Gly Pro ٧ Lys Pro Tyr 돳 A O Pro Leu Lys Gly Asp Gly 90 Trp Asp Cys Thr G S 부 116 Arg 교 쟉 ۷ م Ala Ser Pro Pro Arg Asn (I e tro Glu 118 Gly Pro GIV Gly Ser SSO SIZ 9.90 5.70 **22** 510 130 870 Gin Se. 23 ₹∓ A550 250 5er 52 \$33 \$430 Lys Gin <u>0</u> GIZ Phe Asp Lys Ser Met 11e Arg Ŧ Ser 잣 Ş Met Çs <u>G</u> Phe Ala Leu Cys Ě Ser Pro HIS Pro Arg Arg Š . 10 cys I . บ เบ I e Asn <u>@</u> Ser Glu Pro Gly Arg È Gly . જે Τχ Asn Gly 돳 Oln Pro ςλs . Sys Glu Lys HIS Pro HIS 흔 Gly Ţ Ala GIn Thr G Y <u>k</u>ø GIn Asp Gin Lys Tyr Trp. <u>a</u> Thr Arg Εİ Pro Gin S C 후 Lys. Asp Κa Leu Gin Ser Asn Gly Leu Gly \$ Cys Leu Gly Asn g J 본 Arg S S Ī 투 ¥ Cys 보 Υa Ϋ́ Ser Asn Pro 다 만 Cys Gin **Tyr** GIn Asp Leu Ser Ser Cys Thr Cys Asn Ser Cys Thr **₽** GIY <u>k</u> Leu Asn Asn ה Asp פֿ ار ار 벁 Ė Arg Ş Ş Ser 늄 Ser Ţ Asp Ty \$ Ŧ Ş Asp ΘŊ <u>k</u> Εľ Asp Asp Asn Asn GIY

Çs

Glu 11e

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Gin Lys Phe Gly

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660 Vai 170 170 170 170 170 649 820 Trp 200 Ser Gly GIn 투 Phe Ser Ty. Gly <u>8</u> Asp Leu Ϋ́ Pro <u>G</u> Ļ g Z Asn Gly HIS Asn Ser Lys Asn ۷a 井 Š]e Ser Ser Val 井 δ Leu Asp Leu <u>8</u> Ser ζ II e Pro 丰 ΤŢ Ser 井 Ν Cys G J . Gln lle Leu Arg Trp Arg Ţ 턴 Fro Val Tyr Ser Ser Ser Asp Lys Ser 井 Asn Ser Ser Ile GIn Thr Pro Ser Pro GIN Tyr Asp Thr Š 잣 7 Ely Arg IIe Leu Ser Asn Ė 井 \$ Ala Ala Asp <u>I</u>e HIS Leu Ser Gln Pro Leu Gln GIn Asp Ν = Ser <u>ე</u> Pro <u>8</u> ۷a Gly Ĕ 69 693 650 Leu 770 Leu 730 Asp 750 Leu <u>6</u>6 ₹ % 28 0-8 670 Ser Pro Asp Asn Leu Pro Glu Po Τρ Gly 띰 민 Pro Ļ ñ 阜 GΥ Pro Asp Leu Ser Şα. GI 투 Ser Ser Ser <u>√</u> ςλs Glu Ale Thr 11e Asp 7, Phe Thr Thr Glu Glu Glu Gly Glu Thr Phe GIU. 투 IIe Va I Lys Trp Lys Cys Asp Pro Val HIS Pro Тrр ٧ Trp Glu Ser Gy Τ̈́ζ Asn 11e Asp Phe Glu Leu Asp Thr Glu Leu Ser 부 Arg Phe <u>9</u> Phe <u>s</u> 井 Ser <u>√</u> G Trp Lys Ser <u>ال</u> <u>6</u> Ser Ser Pro Gly Ţ Ser Asp Val Arg n U 11e Pro 뀰 G S Ala 井 Ala Gly Arg Lys <u>-6</u> ē

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7.00 Va. -D -E Ser <u>8</u> . کا <u>8</u> . ∑e Lys Ale ΛøΙ Ser GIn 보 <u>l</u>e Asn Lys Τ̈́ G Ser Thr Pro Pro Arg Thr Ser . ⊡ Asp Ţ Ala 부 <u>1</u>le <u>></u> <u>k</u> Pro Asp Thr Met Arg Val Thr Trp ۷sb ۸a Arg Val Val Ser Val Phe Thr Thr Leu GIn Pro lle Val 11e Thr Trp GIU Ala Val Š His Ser GIY Asn Pro Тyr Pro Ser Leu Pro 11e Val Pro Ile Ser Asp Thr Ile Ile Pro Pro Val GIN GIY GIY Asn Val Gly Thr Gly Gly Leu Thr 1130 GIn Glu Arg Asp Ala Asn Leu His Leu Glu Ala **№** Val Val 9 Glu Tyr 불 Asn Pro Asp Ile Glu Thr Thr Arg Pro Ser 1190 Asn Ser Leu 1110 Ser -S Pro Gly Thr Thr Gln Pro Arg Ser Pro Lys Ala Thr Gly <u>s</u> Val Gly Asn Leu Gin Pro Ala ౼ Phe Asp Asn Leu Ser Ser 11e <u>√</u>8 Val Leu Arg Asp Ser Gin Gin Gly Gly Phe Lys Leu Gly Pro Thr Arg GIC Arg Arg Gly Pro Lys Glu Ser Trp Glu Leu Arg Phe Thr G S Asn Thr Thr Asn Gly Asp Ser Glu Ser <u>G</u> Ser Τχ Leu Thr Leu Arg Thr Val <u>|</u> Ser Pro ה Arg 귤

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Fig. 5D

1560 Gly Arg Ile Ala Na Pro Val Thr 부 Lys Glu Ser Pro Val Lys Asn Glu Glu Asp Val Leu Thr Asn Leu Leu Pro ile Thr 8 Ser Met Gin Val Ν Ŧ Asp . Per Val Asp Ser Lys Thr Val Se Ser Val Leu Lys Pro Ala ۷a Pro Asp ξ Leo 부 Asn ۸a Tyr Ile Ser <u>e</u> 부 Ala Leu Lys Asp Ser Ser Pro Gly Ser Ser 투 Ser Gly Ser Va. Gln Lys Asp <u>5</u> Š G S Set Asp Lys Pro Ser Ser Pro Thr Val 누 <u>8</u> Leu lle ۲ø۱ 井 Arg ' 부 Asp <u>G</u> 부 GIN HIS Glu Thr Ser]e Thr Ile Leu Val <u>8</u> Ser Pro . 부 1530 Lys Trp Leu Pro Pro <u>:</u> Leu Arg Val Ser Ť Asp Ŋ Ala Gln Gln Pro Arg . Je ۷ Gin Gly 1550 Gly Pro Gly Pro Asp Gly Ser g S Ser 8 gr PO Ser Thr 1510 Glu 11e 1490 Val Thr The Ţ Tyr. 1430 Pro Glu Ile Asn Leu Ala Lys Asn פות Phe Olc. Asp ارجح. Arg Pro Arg Ile Ala 본 Ser Val Leu Thr Glu Val Na Leu Leu È Ser Ser Asn Phe Leu Val Lys 116 Arg Gly Ϋ́ Š Asp 본 Ala Ala Ser Ser Pro Leu <u>6</u> = . Pg Pro 보 Pro Leu Ţ Gly Val ķ Ser Ser Λa Asp Met Lys Val H H Asn Ser 부 Asn ş Pro Ser Ser Asn Val Ser ट ट्रे Ile Asn ٦ 일 Asn Val Arg Ile S D ۷ Leu Thr 부 ۷al Asp Met 부 o S 부 ٦ Olc. 2 Ş 뵨 Se ιζs à Pro Phe ۷a Asp J D 표 Š 후]e Arg Leu **1**7 eu Met <u>S</u> Ύβ ٦ Ϊ Asp ٨ G Asp Pro 116 Ser 얼 Asp \<u>a</u> Arg Asn Ser Asn 보

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Fig. 5E

1860 178 1880 170 1980 Ser 2020 Leu Tyr Lys Ile 2000 Thr Trp Cys His Asp Asn Gly Leu Pro Pro Ang Ang Ala Thr Lys Thr Glu Thr ᅣ Asn Ser Leu Leu Pro Ser Νa ZIS Ser ᅽ = 듄 Ser Lys Leu Leu Cys GIn 井 <u>8</u> Pro Pro Asn Val Arg Ą Lys Ile Asp Ala 141 Asn 본 Ser Ser Pro Gly Phe Arg lle Gln Thr Asp Asn Tyr Glu Glu Tyr Gly Gly Asn AB Val Va] Thr Val S Lys Pro ģ GIN Leu Pro Gly Val Pro Leu 부 Leu Ţ Val]]e Τ̈́Υ Val S C Ser IIe IIe Lys Ŧ n L Asn Phe Pro . Б HIS Gin Phe Arg Thr Ser Ile Ser Trp Arg <u>s</u> 부 Pro 투 ٧ø Ala Ser Pro Arg Phe Gln Aso Thr Gin Gn Pro Pro Arg Glu Asn Val 11e Leu] |-<u>olo</u> 첫 √a Va Ę G √ Phe Asp Gly Ser Len Asn Gly Pro Gly Pro Phe Glu Thr Ala Ţ Λa ۷ Aso Glu Glu Val Ser 1950 HIS Arg F Asn (Asp Ala ٥١ Arg Asp 1990 Pro 1730 Ala 950 11e Pro Arg 딍 Pro Arg Asp Asp 부 Lys Lys 1 e Met Pro 11e Arg Thr Arg Thr 11e Lau Arg 부 Arg Met Phe Arg Asn Ala Trp Ala Arg g L S C ۷a 후 ξ S G Asp √a_ 후 Val 보 Ala Asp Asn Ala Ν J G Arg Pro Ser Asp Len Lys Trp Glu Gly His Ŗ H_S S L Asp Asn 뉴 Thr GI√ GIn Pro Val Ser Arg G Z G Z Ile Ser Gly Ş G S 부 Gly HIS Arg G S Ala 井 Glu 누 æ Ŧ Arg Pro Pro <u>G</u> Š \$ Leu 11e G L Lea P 0 Gin Arg ۷aI Leu Asn Asp Ala <u>Б</u> Ser 투 Š Ĕ Phe Asp Pro D D Pro Asn <u>ulo</u> Pro Gin Leo Asp Trp GIn Pro Τζ Pro Pro ΗS <u>ق</u> 부 Pro Leu <u>A</u> Asp <u>8</u> Ser Val ፠ Ja, Leu <u>A</u> Ser Ser

Cys Thr Cys Leu Gly Asn Gly Lys Gly Glu Phe Lys Cys Asp Pro His Glu Ala Thr Cys 2140

Tyr Asp Asp Gly Lys Thr Tyr His Val Gly Glu Gln Trp Gln Lys Glu Tyr Leu Gly Ala

Ile Cys Ser Cys Thr Cys Phe Gly Gly Gly Arg Gly Trp Arg Cys Asp Asn Cys Arg Arg

Pro Gly Gly Glu Pro Ser Pro Glu Gly Thr Thr Gly Gln Ser Tyr Asn Gln Tyr Ser Gln

Arg Tyr His Gln Arg Thr Asn Thr Asn Val Asn Cys Pro Ile Glu Cys Phe Met Pro Leu

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Asp Val Gln Ala Asp Arg Glu Asp Ser Arg Glu GIU ASh GIY GIN Met Met Arg Gin Gly GIU Lys Trp Asp Val Asn Tyr Lys Ile Gly

Fig. 5F

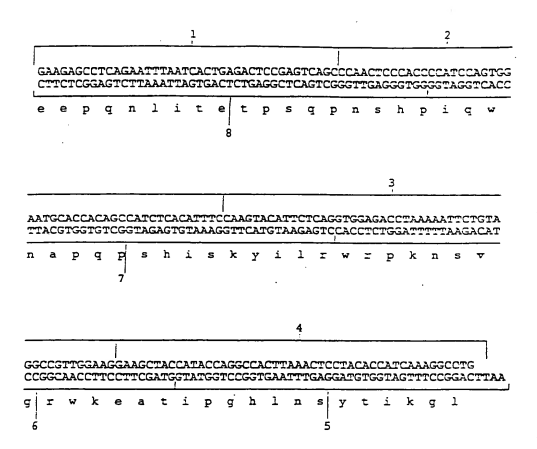


Figure 6 Linker 5 showing the eight constituent oligonucleotides

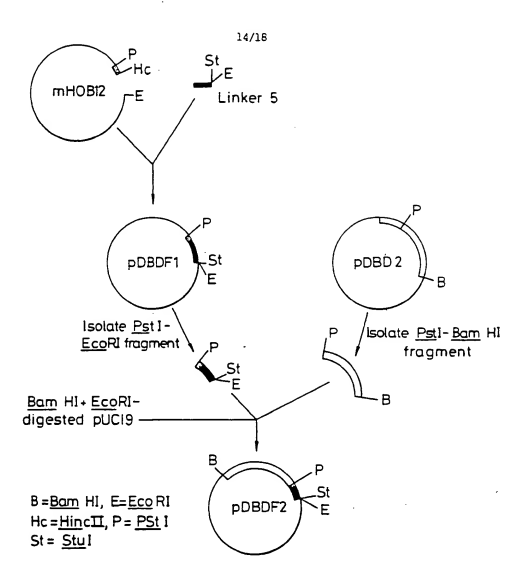


Fig. 7 Construction of pDBDF2

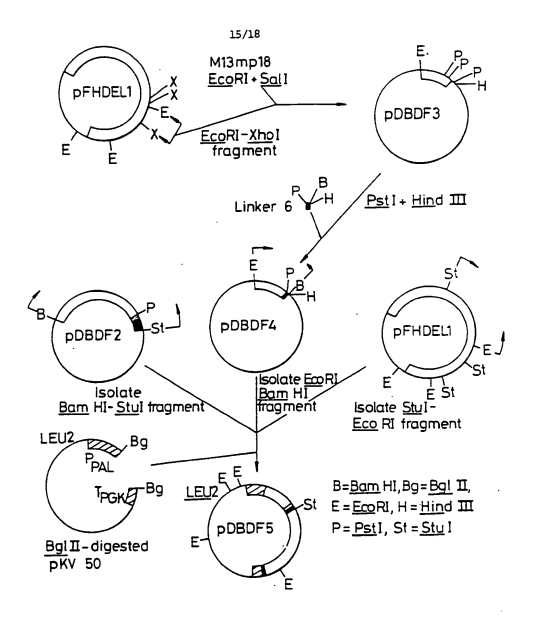


Fig. 8 Construction of pDBDF5

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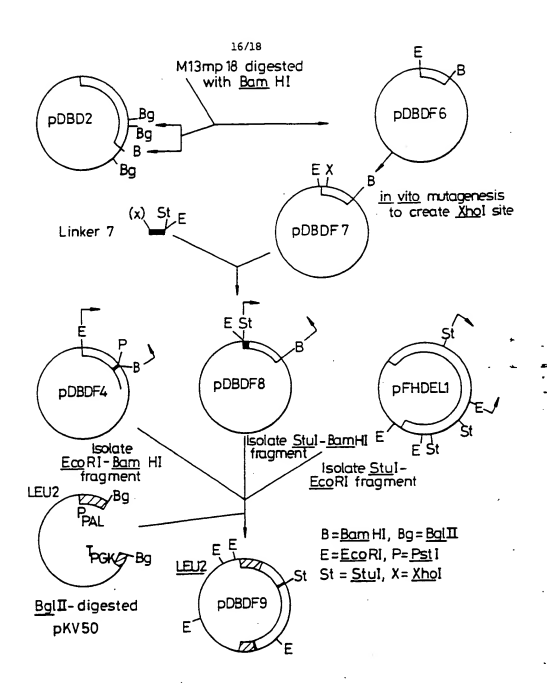
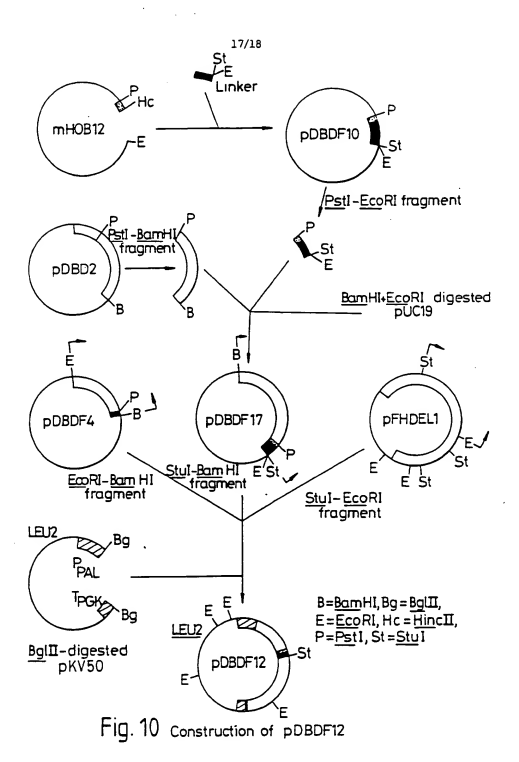


Fig. 9 Construction of pDBDF9

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Figure 11

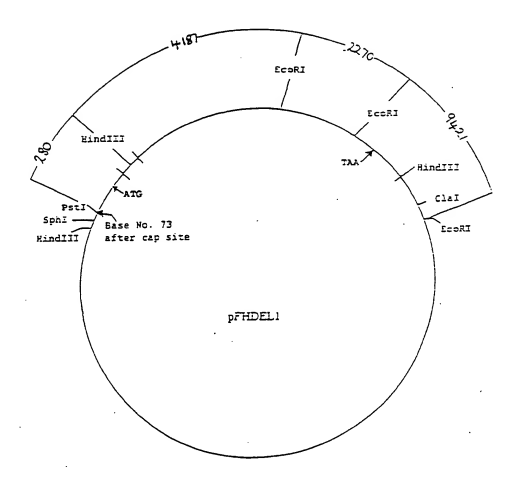
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Name:

pFHDEL1

Vector: pUC18 Amp^{fy} 2860bp

Insert: hFNcDNA = 7630bp



INTERNATIONAL SEARCH REPORT

International Application No PCT/GB 90/00650 I. CLASSIFICATION OF SUBJECT MATTER (II several classification sympols apply, indicate all) 1 According to International Patent Classification (IPC) or to both National Classification and IPC IPC⁵: C 12 N 15/62, C 07 K 13/00, C 12 P 21/02 IL FIELDS SEARCHED Minimum Documentation Searched 7 Classification System i Classification Symbols IPC⁵ C 12 N, C 12 P, C 07 K Documentation Searched other than Minimum Documentation to the Extent that such Documents are included in the Fields Searched * III. DOCUMENTS CONSIDERED TO BE RELEVANT Category * ; Citation of Document, " with indication, where appropriate, of the relevant passages 12 | Relevant to Claim No. 13 Α EP, A, 0308381 (SKANDIGEN et al.) 22 March 1989 EP, A, 0322094 (DELTA BIOTECHNOLOGY LTD)
28 June 1989 T (cited in the application) -----* Special categories of cited docum tater document published after the international filing date or promy date and not in conflict with the application but clied to understand the principle or theory underlying the investion. "A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier document but published on er after the international filing date "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step "L" document which may throw doubts on priority claim(s) or which is clied to establish the publication date of enother casbon or other special reason (as specified) Socument of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person shilled in the art. "O" document reterring to an orei disclosura, was, exhibition or other means "P" document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family IV. CERTIFICATION Date of the Actual Completion of the International Search Date of Mailing of this International Search Report 09.08.90 10th July 1990 M. SOTELO International Searching Authority Signature of Aythonizes Officer EUROPEAN PATENT OFFICE

Form PCT/ISA/210 (second sheet) (January 1885)

ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

GB 9000650

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This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 31/07/90

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Patent document cited in search report EP-A- 0308381	Publication date 22-03-89	Patent family member(s)		Publication date
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